Structure of RGL3 and alignment of RasGEFN,

and RA motifs. RasGEF

R. RASGEF RasGEFN

RGL3 (710 a.a.)

FIG. 1A

212

150

140

749 DHHFYDFEKDPMLLEKLLN-FLEHVNGK-SMRKWVDSVLKIVQRKN 792 168 ASHPEDFGSEVKGQLDRLE-SFLLRTGYAAREGVVGGSADLIRNLR

gi 1354501

gi 158471 544403

gi

consensus

RGL3

194 DQYSEDFCQP-PDFPCLKQlVAYVQLNM-PGSDLERRAHLLLAQLE

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RasGEFN motif	tif		
		10 20 30 40 .50 60	
consensus	Н	CKGGLIKGGTLEKLIEHLTEARDK-VDPTFVETFLLTYRSFITTQELLQKLLYRYNAIPP 5	59
RGL3	64	SKVRVLRAARLERLVGELVFGDRE-QDPSFMPAFLATYRTFVPTACLLGFLLPPMPPP	120
gi 1354501	87	RSSRRLRAGTLEALVRHLLDARTagADMMFTPALLATHRAFTSTPALFGLVADRLEALES	146
gi 158471	636	AGVPMIKGATLCKLIERLTYHIYADPTFVRTFLTTYRYFCSPQQLLQLLVERFNIPDP	693
gi 544403	111	CKVRTVKAGTLEKLVEHLVPAFQ9-SDLSYVTVFLCTYRAFTTTQQVLDLLFKRYGCILP	169
		001 001 00	
		· · · · · · · · · · · · · · · · · · ·	
consensus	9	EGVENRVLNILRLWV	88
RGL3	121	PPPGVVVEIKKTAVQDLSFNknlRAVVSVLGSWL	152
gi 1354501	147	YPPGELERTTGVAISVLSTWL	167
gi 158471	694	SLVyqdtgtagaggmggvggDkehknshredwkRYRKEYVQPPRVLNVLRHWV	748
gi 544403	170	YssRPQDQLKNAISSILGTWL	193

10 20 30 40 50 60	LILLIDPKELAEQLTLIDFELFRKIDPSELLGSVWGKRSKKSPSPLNLERFIERFNE 56 LLDFSVDEVAEQLTLIDLELFSKVRLYECLGSVWSQRDRPGAAGASPTVRATVAQFNT 300 LLTLHPIEIARQLTLLESDLYRAVQPSELVGSVWTKEDKEINSPNLLKMIRHTTN 225 LLLFPPDLVAEQFTLMDAELFKKVVPYHCLGSIWSQRAKKGKEHlapTIRATVAQFNN 420 VLVFLADHLAEQLTLLDAELFLNLIPSQCLGGLWGHRDRPGHSHlcpSVRATVTQFNK 296	70 80 100 110 120 120	130 140 150 160 170 180 * * * * * VSALSSSPISRLKKTWEKLPSKYKKLFEELEELLDPSRNFKNYREALSSCN 151 LSALQSNPIYRLKRSWGAVSREPLSTFRKLSQIFSDENNHLSSREILFQEEategsqeed 406 VSAMNSSPVYRLDHTFEQIPSRQKKILEEAHELSEDHYKKYLAKLRSIN 318 LSALQSNAIHRLKKTWEEVSRGSFRVFQKLSEIFSdenNYSLSRELLIKEGtskfatlem 526 VSALQSSPIHRLRAAWGEATRDSLRVFSSLCQIFSeeDNYSQSRELLVQEVKlqspleph 414	190 210 220 230 240*
1C motif	1 243 171 363 6 239	57 301 226 421 6 297	101 347 270 467 6 355	152 407 319 527 6 415
FIG. Rasgef m	consensus RGL3 1BKD_S gi 544403	consensus RGL3 1BKD_S gi 544403	consensus RGL3 1BKD_S gi 544403 gi 691995	consensus RGL3 1BKD_S gi 544403 gi 691995

1 Մ	4.40
FIG.	100 C C C

(Continued)	q	250 260	270	280	290	
		* · · · · · · · · · · · · · · · · · · ·	*:	*:	*:	
consensus	195	195 REIRQLQSQPYNLRPNRSDIQSLLQQSLDSLPEENELYELSLRIEPRV 242	LQQSLDSLP	EENELYE	LSLRIEPRV	242
RGL3	460	460 ARIQQLQRrcQSYTLSPHPPILAALHAQNQLTEEQSYRLSRVIEPPA 506	HAQNQLT	EEQSYR	LSRVIEPPa	506
1BKD_S	364	364 GEIQQYQNQPYCLRVE-SDIKRFFENLnpMGNSMEKEftdYLFNKSLEIEPRN 415	FENLnpMGNSM	EKEftdYLFN	KSLEIEPRN	415
gi 544403	584	gi 544403 584 AQIKLLQSacNNYSIVPE-EHFGAWFRAMGRLSEAESYNLSCELEPPS 630	FRAMGRLS	EAESYN	LSCELEPPS	630
qi 6919956	468	qi 6919956 468 SELRRLONecRGYNLOPD-HDIORWLOGLRPLTEAOSHRVSCEVEPPG 514	LOGLRPLT	EAOSHR	VSCEVEPPG	514

		09	671	75	63	73	
09 .	*	YALVEVLSGDK	YQLFQVLPGDR	YELVQVISEDK	FELVQLLPGDR	YELLQIISEDH	
. 50	*	RLDDEDPEE	NVPQPWACD	NLESDPAEE	NRDSAVASEI	NLDEDEPED	
40	*	VVQLALEKE	VVRRALQKH	VIQRAMSKH	VISRVLKKN	'VIRKAMDKH	
30	· · · · · · · · · · · · · · · · · · ·	VSSEDTAPD	LTSQDKAPS	1LTSQDKTPA	VTSQDKAPS	JVTSQDKAPT	
50	*:	KPGVAYKTIF	H-GNLYRSII	-NGNMYKSIN	eDGSVYKSII	-NGNMYKSII	
10		1 DQGVLRVYFQDLKPGVAYKTIRVSSEDTAPDVVQLALEKFRLDDEDPEEYALVEVLSGDK 60	613 earvirvsidndh-Gnlyrsilltsqdkapsvvrralqkhnvpqpwacdyqlfqvlpgdr 671	17 DTCIIRISVEDn-NGNMYKSIMLTSQDKTPAVIQRAMSKHNLESDPAEEYELVQVISEDK 75	4 DCRIIRVQMELgeDGSVYKSILVTSQDKAPSVISRVLKKNNRDSAVASEFELVQLLPGDR 63	15 DCCIIRVSLDVd-NGNMYKSILVTSQDKAPTVIRKAMDKHNLDEDEPEDYELLQIISEDH 73	
10			613	17	4	15	
FIG RA motif		consensus	RGL3	1EF5_A	1RLF	1LXD_A	

		O1	66)3	0	00
		92	9	1	9	Ä
06	: : : : :	LSLRFLLKRR	APRDFMLRRKE	FDFILRKKN	HDFLLRQRR	YDFILKKR-
80	***************************************	61 ERKLPDDENPLQLRLNLPRDGLSLRFLLKRRD	672 VLLIPDNANVFYAMSPVAPRDFMLRRKE 699	76 ELVIPDSANVFYAMNSQVNFDFILRKKN 103	64 ELTIPHSANVFYAMDGASHDFLLRQRR	74 KLKIPENANVFYAMNSAANYDFILKKR- 100
70	*:	ERKLPDDENE	VLLI PDNANY	ELVIPDSAN	ELTIPHSAN	KLKI PENAN
		61	672	92	64	74
		consensus	RGL3	1EF5_A	1RLF	1LXD A

Structure of the RGL3 gene (Chr. 19p13.2)

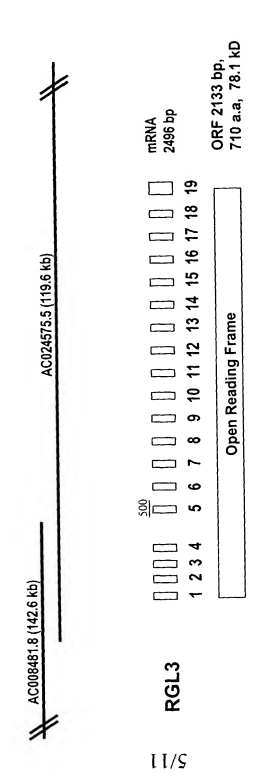


FIG. 2

RGL3

nt: SEQ ID NO: 1 aa: SEQ ID NO: 3

CACT	rgag <i>a</i>	\GGG#	\CGGG	GCGCC	CAGCO							K AAA	7 45
E GAG	L CTG	A GCC	L CTG	A GCA	P CCG	L CTG	Q CAG	D GAC	W TGG	G GGT	E GAA	E GAG	20 84
	E GAG											R CGG	33 123
	R CGC											G GGG	46 162
G GGC	S AGC	Q CAG	A GCT	P CCC	S AGC	P CCC	I ATT	A GCC	N AAT	T ACC	F TTC	L CTC	59 201
	Y TAT												72 240
R CGC	L CTG	E GAG	R CGG	L CTG	V GTG	G GGA	E GAG	L TTG	V GTG	F TTT	G GGA	D GAC	85 279
	E GAG												98 318
	Y TAC												111 357
	L CTG											G GGG	124 396
												F TTC	

FIG. 3

												S TCC	
									R CGA				163 513
									T ACC			G GGC	176 552
									Q CAA				189 591
									A GCT				202 630
									T ACA				215 669
									S TCT			A GCC	228 708
C TGC									P CCT				241 747
									V GTG			Q CAG	254 786
									S TCC				267 825
									S TCG			D GAC	280 864
									T ACT				293 903
												L CTG	306 942
G GGT									A GCC			Q CAG	
R AGG	A GCG	Q CAG	R CGG	L CTG	E GAG	K AAG	W TGG	I ATC	R CGC	I ATC	A GCC	Q CAG	332 1020

FIG. 3

R CGC	C TGC	R CGA	E GAA	L CTG	R CGG	N AAC	F TTC	S TCC	S TCC	L TTG	R CGC	A GCC	345 1059
I ATC	L CTG	S TCC	A GCC	L CTG	Q CAA	s TCT	N AAC	P CCC	I ATC	Y TAC	R CGG	L CTC	358 1098
K AAG	R CGC	S AGC	W TGG	G GGG	A GCA	V GTG	S AGC	R CGG	E GAA	P CCG	L CTA	S TCT	371 1137
T ACT	F TTC	R AGG	K AAA	L CTT	S TCG	Q CAG	I ATT	F TTC	S TCC	D GAT	E GAG	N AAC	384 1176
N AAC	H CAC	L CTC				E GAG						E GAG	397 1215
												G GGC	410 1254
S AGC		P CCC	S TCA	K AAA	P CCA	P	P CCA	G GGC	P CCT	V GTC	P CCC	Y TAC	423 1293
L CTT						D GAC						T ACA	
A GCC	L CTG	P CCG	D GAT	M ATG	L TTG	E GAG	G GGG	D GAT	L CTC	I ATT	N AAC	F TTT	449 1371
						W TGG						I ATC	
Q CAG	Q CAG	L CTG	Q CAG	R AGG	R CGC	C TGT	Q CAG	S AGC	Y TAC	T ACC		S AGC	475 1449
P CCC	H CAC											N AAC	488 1488
Q	L	T	E	E	Q	S	Y	R	L	S		V	
CAG	CTC	ACC	GAG	GAG	CAG	AGC	IAC	CGG	CIC	100	CGG	010	132,

FIG. 3

Т	R	R	R	I	s	L	Т	K	R	L	s	Α	527
ATC	CGA	CGG	CGG	ATC	AGC	CTC	ACC	AAG	CGT	CTC	AGT	GCG	1605
к	L	Α	R	E	K	s	S	S	P	S	G	S	540
AAG	CTT	GCC	CGA	GAG	AAA	AGC	TCA	TCA	CCT	AGT	GGG	AGT	1644
p	G	D	Р	s	S	P	T	S	S	V	S	P	553
CCC	GGG	GAC	CCC	TCA	TCC	CCC	ACC	TCC	AGT	GTG	TCC	CCA	1683
G	S	P	P	S	s	P	R'	S	R	D	Α	P	566
GGG	TCA	CCC	CCC	TCA	AGT	CCT	AGA	AGC	AGA	GAT	GCT	CCT	1722
Ά	G	S	P	Р	Α	S	P	G	P	Q	G	P	579
GCT	GGC	AGT	CCC	CCG	GCC	TCT	CCA	GGG	CCC	CAG	GGC	CCC	1761
S	т	K	L	P	L	s	L	D	L	P	S	P	592
AGC	ACC	AAG	CTG	CCC	CTG	AGC	CTG	GAC	CTG	CCC	AGC	CCC	1800
R	P	F	Α	L	P	L	G	S	P	R	I	P	605
CGG	CCC	TTC	GCT	TTG	CCT	CTG	GGC	AGC	CCT	CGA	ATC	CCC	1839
L	P	Α	Q	Q	S	S	E	Α	R	V	1	R	618
CTC	CCG	GCG	CAG	CAG	AGC	TCG	GAG	GCC	CGT	GTC	ATC	CGC	1878
V	S	I	D	N	D	Н	G	N	L	Y	R	S	631
GTC	AGC	ATC	GAC	AAT	GAC	CAC	GGG	AAC	CTG	TAT	CGA	AGC	1917
I	L	L	${f T}$	S	Q	D	K	Α	P	S	V	V	644
ATC	TTG	CTG	ACC	AGT	CAG	GAC	AAA	GCC	CCC	AGC	GTG	GTC	1956
R	R	Α	L	Q	K	H	N	V	P	Q	P	W	657
CGG	CGA	GCC	TTG	CAG	AAG	CAC	AAT	GTG	CCC	CAG	CCC	TGG	1995
Α	С	D	Y	Q	L	F	Q	V	L	P	G	D	670
GCC	TGT	GAC	TAT	CAG	CTC	TTT	CAA	GTC	CTT	CCT	GGG	GAC	2034
R	V	L	L	I	P	D	N	Α	N	V	F	Y	683
CGG	GTG	CTC	CTG	ATT	CCT	GAC	AAT	GCC	AAC	GTC	TTC	TAT	2073
Α	М	S	P	V	Α	P	R	D	F	M			696
GCC	ATG	AGI	CCA	GTC	: GCC	CCC	AGA	GAC	TTC	ATG	CTG	CGG	2112
R	к	E	G	т	R	N	Т	L	s	V	S	P	709
CGG	AAA	GAG	GGG	ACC	CGG	AAC	ACT	CTG	TCI	GTC	TCC	CCA	2151

FIG. 3

s *		711
AGC TGA	GGCAGCCCTGTCCTCCACAAGACACAAGTCCCACAGGCAAG	2200
CTTGCGACT	CTTCTCCTGGAAAGCTGCCATCCCCCAGTAGAGGCCACTGTGC	2252
TGTGTATCC	CAGGACCACCCAACTGTAGCCCATTGGACCCCATCTCTTT	2304
TTCTGACTC'	IGTTGGTACTAGATCCATATTCCAAAGACATCAGCCCATGGGT	2356
GGCTGGTGG	AGAGCTCAATCCCATAAATGTAGAAAGAGGTGGGGCATGGATA	2408
CGTCAAATC	CCTCCCCAGAGAAATCTTATAAATGTTAGAGACGCATCAGAAG	2460
TGACAGATG	CGGATGAAAATAGTGACCAGAGTTATG	2496

Tissue Expression profile of RGL3

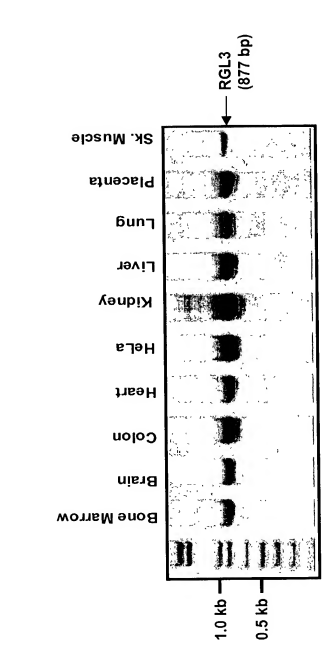


FIG. 4